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DOUBLE CURRENT COLLECTOR NEGATIVE ELECTRODE
DESIGN FOR ALKALI METAL ION ELECTROCHEMICAL CELLS

CROSS REFERENCE TO RELATED APPLICATION

This application claims priority based on provisional application Serial No. 60/249,688, filed
November 17, 2000.

BACKGROUND OF THE INVENTION

This invention relates to the conversion of chemical energy to electrical energy. In particular, the present invention relates to a new negative electrode design having an anode active material sandwiched between two current collectors with an anode material capable of intercalating and de-intercalating the anode active material in contact with the opposite sides of the current collectors. The present negative electrode design is useful for high discharge rate secondary cells, such as those powering an implantable medical device.

SUMMARY OF THE INVENTION

Secondary electrochemical cells are typically built
25 in a discharged state having a negative electrode of a
carbonaceous material and a positive electrode of a
lithiated material. A reverse potential is then applied
to the cell to cause lithium ions at the positive
electrode to migrate through the electrolyte and
30 intercalate into the carbonaceous anode material. The

Graphitic forms of carbon are generally preferred for the anode material because they exhibit acceptable cycling capability and good reversible capacity,

35 especially during initial change and discharge cycles.

cell is then discharged in a normal manner.

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5 While graphite is commonly used, there are other carbonaceous materials which have better reversible capacities. One in particular is hard carbon. This material has excellent cycling characteristics, but suffers from a relatively large initial irreversible capacity.

Therefore, according to the present invention, a secondary cell is constructed having a "sacrificial" piece of alkali metal as a portion of the negative electrode along with an alkali metal intercalation and de-intercalation material, such as a carbonaceous anode material. In the case of hard carbon, the sacrificial alkali metal is preferably a piece of lithium metal and is sized to compensate for the initial irreversible capacity of this anode material. After the cell is activated with an electrolyte, the lithium metal automatically intercalates into the hard carbon anode material. That way, the sacrificial lithium metal compensates for the generally unacceptable irreversible capacity of hard carbon. The superior cycling longevity of hard carbon now provides a secondary cell of extended use beyond that known for conventional secondary cells having only graphitic anode materials.

Accordingly, one object of the present invention is to improve the cycling performance of lithium ion cells by providing a new concept in negative electrode design. Further objects of this invention include providing a cell design for improving the capacity and utilization efficiency of lithium-containing secondary cells.

These and other objects of the present invention
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5 in the art by reference to the following description.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

The electrochemical cell of the present invention is of a secondary, rechargeable chemistry. The cell comprises an anode active metal selected from Groups IA, IIA and IIIB of the Periodic Table of the Elements, including lithium, sodium, potassium, etc.

In conventional secondary electrochemical systems, the anode or negative electrode comprises an anode material capable of intercalating and de-intercalating the anode active material, such as the preferred alkali metal lithium. Typically, the anode material of the negative electrode comprises any of the various forms of carbon (e.g., coke, graphite, acetylene black, carbon black, glassy carbon, etc.) which are capable of reversibly retaining the lithium species. Graphite is particularly preferred in conventional secondary cells. "Hairy carbon" is another particularly preferred conventional material due to its relatively high lithium-retention capacity. "Hairy carbon" is a material described in U.S. Patent No. 5,443,928 to Takeuchi et al., which is assigned to the assignee of the present invention and incorporated herein by reference.

30 However, it is known that graphitic forms of carbon do not possess as high of a reversible capacity as hard carbon. Hard carbon, defined as non-graphitizable carbon material, has two to four times as much reversible capacity as graphite. Hard carbon is 35 typically prepared by firing an appropriate organic

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cursor to temperatures between 700°C and 1,200°C. Hard carbons typically exhibit excellent cyclability and high reversible capacity. The current state of the art, i.e., graphite, is theoretically limited to a capacity of 372 mAh/q, while hard carbons exhibit capacities of 400 mAh/g or more. This means that a secondary cell 10 built with hard carbon as the anode material can be charged and discharged, or cycled, two to four times as much as a secondary cell built with a graphitic form of anode material. The improvement in cycle life is based 15 on the dimensional stability of hard carbon during lithium intercalation and de-intercalation. This means that a secondary cell built with hard carbon as the anode material has potentially higher capacity than a secondary cell built with a graphitic form of anode 20 material.

On the other hand, the main reason hard carbon is not used more frequently in secondary cells is because it suffers from large initial cycle irreversible capacity relative to graphite. This irreversible capacity must be compensated for by the addition of cathode active material to the cell which, in turn, lowers the cell's capacity.

For this reason, the negative electrode of the present secondary cells is built in a double screen configuration having a "sacrificial" piece of alkali metal, preferably lithium, sandwiched between two current collectors. A carbonaceous material or a material capable of intercalation and de-intercalation of the alkali metal contacts the opposite side of at least one, and preferably both, of the current

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collectors. The purpose of the sacrificial alkali metal is to compensate for the irreversible capacity of the intercalation and de-intercalation material, for example hard carbon, versus graphite. One embodiment of the present negative electrode has the following configuration:

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hard carbon/current collector/lithium/current collector/hard carbon.

With this double current collector electrode design, the amount of lithium metal is adjusted to precisely compensate for the irreversible capacity of the hard carbon. Upon activating the cell with an ionconductive electrolyte, the alkali metal migrates into the hard carbon resulting in complete consumption of the alkali metal. The absence of the alkali metal in the cell preserves the desirable safety and cycling properties of the intercalation negative and positive electrodes. Therefore, the present negative electrode preferably having lithium metal sandwiched between two current collectors which are further sandwiched between two hard carbon structures has a significantly higher volumetric and/or gravimetric energy density than a conventional negative electrode of a carbonaceous material contacted to a single current collector.

Regardless of the carbonaceous nature or makeup of 30 the anode material, fibers are particularly advantageous. Fibers have excellent mechanical properties which permit them to be fabricated into rigid electrode structures that are capable of withstanding degradation during repeated charge/discharge cycling.

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Moreover, the high surface area of carbon fibers allows for rapid charge/discharge rates.

The carbonaceous portion of the present negative electrode for a secondary cell is fabricated by mixing about 90 to 97 weight percent of the anode material, preferably hard carbon, with about 3 to 10 weight percent of a binder material, which is preferably a fluoro-resin powder such as polytetrafluoroethylene (PTFE), polyvinylidene fluoride (PVDF), polyethylenetetrafluoroethylene (ETFE), polyamides, polyimides, and mixtures thereof.

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This negative electrode admixture is provided on a current collector such as of a copper, stainless steel, titanium, tantalum, platinum, gold, aluminum, nickel, cobalt nickel alloy, highly alloyed ferritic stainless steel containing molybdenum and chromium, and nickel-, chromium-, and molybdenum-containing alloy foil or screen by casting, pressing, rolling or otherwise contacting the admixture thereto.

Another type of anode material useful with the 25 present invention is a metal which can reversibly allow with alkali metals. Such metals include, but are not limited to, Sn, Si, Al, Pb, Zn, and Ag. These alloying metals demonstrate very high reversible capacities, but suffer from poor cycling properties because of massive 30 dimensional changes during the alloying process. One approach which has been used to solve this problem is to suspend nano-sized particles of them in a matrix of inactive material. The inactive material is typically generated by reduction of an oxide of the alloying metal 35 during the first cycle. Examples of oxides include but

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5 are not limited to SnO, SnO₂, SiO, and SnO(B₂O₃)_x(P₂O₅)_y. These oxides exhibit good cycling properties and high reversible capacities, but reduction of the metal oxide consumes large amounts of alkali metal during the first charge. Thus, in conventional secondary chemistries,
10 additional cathode active material must be incorporated into the cell to compensate for this reduction, thereby reducing cell capacity.

It is, therefore, another preferred embodiment of the present invention to place a piece of alkali metal between the above metal oxide anode materials. The amount of alkali metal is chosen to precisely compensate for the reduction of the metal oxide. Then, upon activating the cell with an ion-conductive electrolyte, the alkali metal migrates into the anode material resulting in complete consumption of the alkali metal. As before, the absence of alkali metal in the cell preserves the desirable safety and cycling properties of the anode material, and the resulting cell has capacity which exceeds that of the current art.

Still another preferred embodiment of the present invention takes advantage of active materials that are typically used as cathode active materials in primary cells, but which can not normally be used in conventional secondary cells. The current art in rechargeable cells typically uses the positive electrode as the source of alkali metal ions. This prohibits the use of metal-containing cathode active materials which do not contain alkali metal ions. Examples of such metal-containing materials include V₂O₅, SVO, CSVO, MnO₂, TiS₂, CuO₂, Cu₂S, FeS, FeS₂, CF_x, Ag₂O, Ag₂O₂, CuF,

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5 Ag₂CrO₄, copper oxide, copper vanadium oxide, and mixtures thereof. The use of these active materials in the positive electrode of a secondary cell normally requires the presence of an alkali metal anode or a premetalated anode material (most desirably carbon). The use of an alkali metal anode is undesirable because of the previously discussed poor cycling and safety properties of these materials. Pre-metalated carbon materials are known in the art, but have never been commercialized because of difficulties in reliably
15 manufacturing them.

In the present invention, a piece of alkali metal is used in conjunction with one or more of the previously described metal-containing materials which are typically used as cathode active materials in primary cells. In the present invention, the piece of alkali metal serves as the source of alkali metal ions and is sandwiched between two lavers of one or more of the above metal-containing materials now serving as an anode material. The amount of alkali metal is chosen such that it precisely compensates for the reversible capacity of the anode material. Upon activating the cell with an ion-conductive electrolyte, the alkali metal migrates into the anode material resulting in complete consumption of the alkali metal, which preserves the desirable safety and cycling properties of the anode material.

Now, it is possible for the positive electrode to have as its active material any of the above materials which are described as useful anode materials for the negative electrode of the present invention and which

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5 form an acceptable electrochemical potential with the anode material. As previously described, these materials are not typically used as cathode active materials in secondary cells because they are not lithiated. An exemplary couple would be graphite/Li as 10 a negative electrode and V₂O₅ or SVO as a positive electrode.

As is the case with the above described carbonaceous anode materials, the alloying anode metals and the metal-containing anode materials are formed into a sandwich electrode body for incorporation into an electrochemical cell by mixing one or more of them with one of the above described binder materials. Further, up to about 10 weight percent of a conductive diluent is preferably added to the mixture to improve conductivity. Suitable materials for this purpose include acetylene black, carbon black and/or graphite or a metallic powder such as powdered nickel, aluminum, titanium and stainless steel. The preferred anode material mixture thus includes a powdered fluoro-polymer binder present at about 1 to 5 weight percent, a conductive diluent present at about 1 to 5 weight percent and about 90 to 98 weight percent of the anode material.

Therefore, one exemplary negative electrode has an anode material short circuited to the alkali metal anode active material by parallel connection through the current collectors. The following configuration is exemplary:

first anode material/current collector/alkali metal/current collector/second anode material, wherein the first and second anode materials are either the same 5 or different.

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Another embodiment of the present invention has the alkali metal sandwiched between the anode material, in which the anode material is short circuited to the alkali metal by direct contact. This negative electrode has the configuration:

first anode material/current collector/second anode material/alkali metal/third anode material/current collector/fourth anode material, wherein the first, second, third and fourth anode materials are either the same or different.

A third exemplary embodiment of the present invention has the configuration:

 ${\it anode material/current collector/alkali\ metal,}$ wherein the anode material faces the positive electrode.

More preferred embodiments include the following negative electrode configurations:

hard carbon/current collector/lithium/current collector/hard carbon, or

anode material/current

25 collector/lithium/current collector/anode material, wherein the anode material is selected from the group consisting of SnO, SnO₂, SiO, SnO(B₂O₃)_x(P₂O₅)_y, V₂O₅, SVO, CSVO, MnO₂, TiS₂, CuO₂, Cu₂S, FeS, FeS₂, CF_x, Ag₂O, Ag₂O₂, CuF, Ag₂CrO₄, copper oxide, copper vanadium oxide, and 30 mixtures thereof, or

carbonaceous material/current collector/lithium/current collector/carbonaceous material, or

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graphite/current
collector/graphite/lithium/graphite/current
collector/graphite.

In a secondary cell, the reaction at the positive electrode involves conversion of ions which migrate from the negative electrode to the positive electrode into atomic or molecular forms. The positive electrode preferably comprises lithiated oxides, sulfides, selenides, and tellurides of such metals as vanadium, titanium, chromium, copper, molybdenum, niobium, iron, nickel, cobalt and manganese. Preferred lithiated oxides include $\text{Li}_x \text{Ti}_5 \text{O}_{12}$ (x = 4 to 7), $\text{Li}_{3-x} \text{M}_x \text{N}$ (M = Co, Ni; x = 0.1 to 0.6), LiNiO₂, LiMn₂O₄, LiMnO₂, LiV₂O₅, LiCoO2, LiCo0.92Sn0.08O2 and LiCo1-xNixO2. The abovedescribed alkali metal intercalation and deintercalation materials such as SVO, CSVO, Ag2O, Ag2O2, CuF2, Aq2CrO4, MnO2, V2O5, TiS2, Cu2S, FeS, FeS2, copper oxide, copper vanadium oxide, and mixtures thereof, are also useful as cathode active materials.

To charge such secondary cells, the lithium ions

comprising the positive electrode are intercalated into
the carbonaceous anode material or the lithium
intercalation and de-intercalation anode material of the
negative electrode by applying an externally generated
electrical potential to the cell. The applied

recharging potential draws the lithium ions from the
cathode active material, through the electrolyte and
into the anode material to saturate it. In the case of
carbon, the resulting Li_xC₆ material can have an x
ranging between 0.1 and 1.0. The cell is then provided

with an electrical potential and discharged in a normal

5 manner.

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The above described cathode active materials are formed into a positive electrode by mixing them with one or more of the above described binder materials and conductive diluents. The preferred cathode active mixture thus includes a powdered fluoro-polymer binder present at about 1 to 5 weight percent, a conductive diluent present at about 1 to 5 weight percent and about 90 to 98 weight percent of the cathode active material.

Positive electrodes for incorporation into an electrochemical cell according to the present invention may be prepared by rolling, spreading or pressing the cathode active formulations onto a suitable current collector of any one of the previously described materials suitable for the negative electrode. The preferred current collector material is aluminum. Positive electrodes prepared as described above may be in the form of one or more plates operatively associated with at least one or more plates of a negative electrode, or in the form of a strip wound with a corresponding strip of the negative electrode in a structure similar to a "jellyroll".

In order to prevent internal short circuit conditions, the negative electrode is separated from the positive electrode by a suitable separator material. The separator is of electrically insulative material, and the separator material also is chemically unreactive with the anode and cathode active materials and both chemically unreactive with and insoluble in the electrolyte. In addition, the separator material has a degree of porosity sufficient to allow flow there

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through of the electrolyte during the electrochemical reaction of the cell. Illustrative separator materials include fabrics woven from fluoropolymeric fibers including polyvinylidine fluoride,

polyethylenetetrafluoroethylene, and

polyethylenechlorotrifluoroethylene used either alone or laminated with a fluoropolymeric microporous film, non-woven glass, polypropylene, polyethylene, glass fiber materials, ceramics, polytetrafluoroethylene membrane commercially available under the designation 15 ZITEX (Chemplast Inc.), polypropylene membrane commercially available under the designation CELGARD (Celanese Plastic Company, Inc.) and a membrane commercially available under the designation DEXIGLAS

(C.H. Dexter, Div., Dexter Corp.).

The electrochemical cell of the present invention further includes a nonaqueous, ionically conductive electrolyte which serves as a medium for migration of ions between the negative and positive electrodes during electrochemical reactions of the cell. A suitable electrolyte has an inorganic, ionically conductive salt dissolved in a nonaqueous solvent, and more preferably, the electrolyte includes an ionizable alkali metal salt

dissolved in a mixture of aprotic organic solvents comprising a low viscosity solvent and a high 30 permittivity solvent. The inorganic, ionically

conductive salt serves as the vehicle for migration of the anode ions to intercalate or react with the cathode active materials. Preferably, the ion forming alkali metal salt is similar to the alkali metal comprising the

35 anode active material.

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In the case of an anode active material comprising lithium, the alkali metal salt of the electrolyte is a lithium based salt. Known lithium salts that are useful as a vehicle for transport of alkali metal ions between the negative electrode and the positive electrode include LiPF₆, LiBF₄, LiAsF₆, LiSbF₆, LiClO₄, LiO₂, LiAlCl₄, LiGaCl₄, LiC(SO₂CF₃)₃, LiN(SO₂CF₃)₂, LiSCN, LiO₃SCF₃, LiC₆F₅SO₃, LiO₂CCF₃, LiB(C₆H₅)₄, LiCF₃SO₃, and mixtures thereof.

Low viscosity solvents useful with the present invention include esters, linear and cyclic ethers and dialkyl carbonates such as tetrahydrofuran (THF), methyl acetate (MA), diglyme, trigylme, tetragylme, dimethyl carbonate (DMC), 1,2-dimethoxyethane (DME), 1,2diethoxyethane (DEE), 1-ethoxy, 2-methoxyethane (EME), ethyl methyl carbonate (EMC), methyl propyl carbonate, ethyl propyl carbonate, diethyl carbonate (DEC), dipropyl carbonate, and mixtures thereof, and high permittivity solvents include cyclic carbonates, cyclic esters and cyclic amides such as propylene carbonate (PC), ethylene carbonate (EC), butylene carbonate, acetonitrile, dimethyl sulfoxide, dimethyl formamide, dimethyl acetamide, y-valerolactone, y-butyrolactone (GBL), N-methyl-pyrrolidinone (NMP), and mixtures thereof.

A preferred electrolyte for a secondary cell according to the present invention comprises a solvent mixture of EC:DMC:EMC:DEC. Most preferred volume percent ranges for the various carbonate solvents include EC in the range of about 20% to about 50%; DMC in the range of about 12% to about 75%; EMC in the range

of about 5% to about 45%; and DEC in the range of about 3% to about 45%. In a preferred form of the present invention, the electrolyte activating the cell is at equilibrium with respect to the ratio of DMC:EMC:DEC. This is important to maintain consistent and reliable cycling characteristics. It is known that due to the 10 presence of low-potential (anode) materials in a charged cell, an un-equilibrated mixture of DMC:DEC in the presence of lithiated graphite (LiC6~0.01 V vs Li/Li+) results in a substantial amount of EMC being formed. 15 When the concentrations of DMC, DEC and EMC change, the cycling characteristics and temperature rating of the cell change. Such unpredictability is unacceptable. This phenomenon is described in detail in U.S. patent application Serial No. 09/669,936, filed September 26, 20 2000, which is assigned to the assignee of the present invention and incorporated herein by reference. Electrolytes containing the quaternary carbonate mixture of the present invention exhibit freezing points below -50°C, and lithium ion secondary cells activated with

25 such mixtures have very good cycling behavior at room temperature as well as very good discharge and charge/discharge cycling behavior at temperatures below -40°C.

The assembly of the secondary cells described

30 herein is preferably in the form of a wound element configuration. That is, the fabricated negative electrode, positive electrode and separator are wound together in a "jellyroll" type configuration or "wound element cell stack" such that the negative electrode is

35 on the outside of the roll to make electrical contact

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5 with the cell case in a case-negative configuration. Using suitable top and bottom insulators, the wound cell stack is inserted into a metallic case of a suitable size dimension. The metallic case may comprise materials such as stainless steel, mild steel,
10 nickel-plated mild steel, titanium, tantalum or aluminum, but not limited thereto, so long as the metallic material is compatible for use with the other cell components.

The cell header comprises a metallic disc-shaped body with a first hole to accommodate a glass-to-metal seal/terminal pin feedthrough and a second hole for electrolyte filling. The glass used is of a corrosion resistant type having up to about 50% by weight silicon such as CABAL 12, TA 23, FUSITE 425 or FUSITE 435. The positive terminal pin feedthrough preferably comprises titanium although molybdenum, aluminum, nickel allov, or stainless steel can also be used. The cell header is typically of a material similar to that of the case. The positive terminal pin supported in the glass-to-metal seal is, in turn, supported by the header, which is welded to the case containing the electrode stack. The cell is thereafter filled with the electrolyte solution described hereinabove and hermetically sealed such as by close-welding a stainless steel ball over the fill hole, but not limited thereto. The above assembly describes a case-negative cell,

The above assembly describes a case-negative cell, which is the preferred construction of the exemplary secondary cell of the present invention. As is well known to those skilled in the art, the present secondary electrochemical systems can also be constructed in case-

5 positive configuration.

It is appreciated that various modifications to the inventive concepts described herein may be apparent to those of ordinary skill in the art without departing from the spirit and scope of the present invention as defined by the appended claims.